

Europäisches Patentamt
European Patent Office
Office européen des brevets



EP 0 737 222 B1

(12)

# **EUROPEAN PATENT SPECIFICATION**

- (45) Date of publication and mention of the grant of the patent: 30.09.1998 Bulletin 1998/40
- (21) Application number: 95905881.9
- (22) Date of filing: 21.12.1994

- (51) Int Cl.<sup>6</sup>: **C08G 67/04**, C09D 163/00, C08L 73/02, C08G 59/68
- (86) International application number: PCT/US94/14166

(11)

- (87) International publication number: WO 95/18173 (06.07.1995 Gazette 1995/29)
- (54) HIGH SOLIDS COATING COMPOSITION

  ÜBERZUGSMITTEL MIT HOHEM FESTSTOFFANTEIL

COMPOSITION DE REVETEMENT A HAUTE TENEUR EN MATIERES SOLIDES

- (84) Designated Contracting States: BE CH DE DK ES FR GB IT LI NL SE
- (30) Priority: 29.12.1993 US 175819
- (43) Date of publication of application: 16.10.1996 Bulletin 1996/42
- (73) Proprietor. E.I. DU PONT DE NEMOURS AND COMPANY
  Wilmington Delaware 19898 (US)
- (72) Inventors:
  - BARSOTTI, Robert, J.
     Franklinville, NJ 08322 (US)
  - JOHNSON, Jeffery, W. Rochester Hills, MI 48307-3147 (US)

- (74) Representative: Balley, David Martin Brookes & Martin High Holborn House • 52-54 High Holborn London WC1V 6SE (GB)
- (56) References cited: EP-A- 0 081 163 WO-A-94/11415 US-A- 1 993 828 US-A- 1 993 828
  - DATABASE WPI Derwent Publications Ltd., London, GB; AN 92-068793(09) & JP,A,4 012 301 (FUJI PHARM IND KK) 16 January 1992

Note: Within nine months from the publication of the mention of the grant of the European patent, any person may give notice to the European Patent Office of opposition to the European patent granted. Notice of opposition shall be filed in a written reasoned statement. It shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention).

### Description

5

The present invention relates to a curable coating composition particularly useful as a top coat in multi-layered

Base coat-clear coat systems have found wide acceptance in the past decade as automotive finishes. Continuing effort has been directed to such coating systems to improve the overall appearance, the clarity of the top coat, and the resistance to deterioration. Further effort has been directed to the development of coating compositions having low

Previous efforts at improving the etch resistance and durability of coatings had suggested the use anhydride resins volatile organic content (VOC). having pendant non-cyclic anhydride moieties in combination with resins that react with the polyanhydride resins to cure under curing conditions. However, a continuing need exists for coating formulations which can be sprayed at exceptionally low VOC and which exhibit outstanding performance characteristics after application, and particularly

The present invention provides a sprayable coating composition which can be easily applied at high solids and resistance to environmental etching. exhibits outstanding appearance and durability after application and ease of maintenance.

Specifically, the instant invention provides a curable coating composition comprising organic solvent and binder comprising (a) an anhydride resin having a molecular weight of less than about 2000 than contains a central moiety, and an average of at least three pendant, terminal, non-cyclic anhydride moieties bonded to each central moiety; (b) an oligomer having epoxy functionality of at least 2 and having a molecular weight of less than about 1500; (c) a functional amount of at least one active tertiary amine catalyst, and (d) about from 0.01 to 5%, by combined weight of the binder component (a) and (b), of a surface tension reducing agent in an amount sufficient to wet the surface onto which it is applied, the composition having a volatile organic contact of less than about 360g/l (3.0 pounds per gallon), and wherein the ratio of equivalents of epoxy to anhydride is about from 0.7 to 1.4.

### DETAILED DESCRIPTION OF THE INVENTION 25

Anhydride resins which can be used in the present invention include those having a molecular weight of less than about 2000 having a central moiety and more than one pendant, non-cyclic anhydride moiety bonded to each central moiety. The anhydride is asymmetrical, and preferably contains a moiety represented by the following formula:

wherein (CM) is a central moiety. (R1) is an organic moiety, and n is a number of pendant anhydride groups that

The central moiety can be a simple organic moiety, such as an aliphatic, cycloaliphatic or aromatic moiety, with a averages greater than one. plurality of anhydride groups bonded to it. Alternatively, it can contain a plurality of repeating units which are bonded to one or more pendant anhydride groups. Examples of suitable non-polymeric central moieties are those derived from multifunctional alcohols such as pentaerythritol, trimethylolpropane and neopentyl glycol. The multifunctional alcohols are reacted with cyclic, monomeric anhydride such as methyl hexahydrophthalic anhydride to give a multifunctional acid containing moiety. The resulting product is then reacted with ketene to form the linear pendant anhydride.

The central moiety is linked to more than one non-cyclic anhydride moiety, on average. It is preferably linked to at least about 2 non-cyclic anhydride groups on average and more preferably to at least about 3 non-cyclic anhydride groups on average. The anhydride equivalent weight (formula weight per anhydride group) is preferably at least about 200 and preferably no more than about 1000.

Each anhydride moiety is typically terminated by an organic group (R1). This group is preferably aliphatic and more preferably alkyl. It preferably contains no more than about 6 carbon atoms, more preferably no more than about 4

The oligomeric anhydride can optionally contain a polyvalent organic moiety (A) that is linked to a plurality of carbon atoms, and most preferably methyl. anhydride groups by a plurality of pendant linking groups (LG), as illustrated in the following formula:

55

30

35

40

5

10

15

20

40

45

50

55

The linking group (LG) can contain, for example, ester linkages, alkylene groups, ether linkages, urethane linkages and combinations of those. The polyvalent organic group can contain, for example, a polyvalent alkyl or aromatic group. The combination of the polyvalent organic moiety (A) and the linking groups (LG) forms the central moiety (CM) as

previously described.

The central moiety can optionally contain other functional groups in addition to the pendant non-cyclic anhydride groups. For example, the central moiety may contain pendant acid groups, so that the anhydride is represented by the formula:

wherein m is the number of pendant acid groups and all other characters have the meaning previously given. The molar ratio of pendant non-cyclic anhydride groups to pendant acid groups in the oligomeric anhydride is preferably at least about 25:75, more preferably at least about 50:50, and more highly preferably at least about 75:25. Most preferably, the anhydride contains substantially no pendant acid groups. The central moiety can also contain minor

quantities of cyclic anhydride moieties.

The molecular weight of the anhydride resin is an important feature of the present invention, and should be less than about 2000. At molecular weights of the oligomeric anhydride greater than 2000, it is difficult to attain a sprayable composition with a volatile organic content of less than about 3.0 pounds of organic solvent per gallon of curable compositions. The molecular weight of the anhydride resin is preferably about from 400 to 1,000, and the anhydride resin preferably has 3 to 4 pendant, non-cyclic anhydride moieties bonded to each central moiety.

The oligomer component contains at least two epoxy groups and should have a molecular weight of less than about 1500. Typical epoxy components are as follows: sorbitol polyglycidyl ether, mannitol polyglycidyl ether, pentaer-ythritol polyglycidyl ether, glycerol polyglycidyl ether, low molecular weight epoxy resins such as epoxy resins of epythritol polyglycidyl ether, glycerol polyglycidyl esters of polycarboxylic acids, polyglycidyl ethers of isocyanurates, ichlorohydrin and bisphenol-A., di- and polyglycidyl esters of polycarboxylic acids, polyglycidyl ethers of isocyanurates, ichlorohydrin and bisphenol-A., di- and polyglycidyl esters of polycarboxylic acids, polyglycidyl ethers of isocyanurates, ichlorohydrin and bisphenol-A., di- and polyglycidyl ether, such as DCE-358® from Dixie Chemical, and disuch as "Denecol" EX301 from Nagase. Sorbitol polyglycidyl ether, such as DCE-358® from Dixie Chemical, and disuch as "Denecol" EX301 from Nagase. Sorbitol polyglycidyl ether, such as DCE-358® from Dixie Chemical, and disuch as "Denecol" EX301 from Nagase. Sorbitol polyglycidyl ether, such as DCE-358® from Dixie Chemical, and disuch as "Denecol" EX301 from Nagase. Sorbitol polyglycidyl ether, such as DCE-358® from Dixie Chemical, and disuch as "Denecol" EX301 from Nagase. Sorbitol polyglycidyl ether, such as DCE-358® from Dixie Chemical, and disuch as "Denecol" EX301 from Nagase. Sorbitol polyglycidyl ether, such as DCE-358® from Dixie Chemical, and disuch as "Denecol" Ex301 from Nagase. Sorbitol polyglycidyl ether, such as DCE-358® from Dixie Chemical, and disuch as "Denecol" Ex301 from Nagase. Sorbitol polyglycidyl ether, such as DCE-358® from Dixie Chemical, and disuch as "Denecol" Ex301 from Nagase. Sorbitol polyglycidyl ether, such as DCE-358® from Dixie Chemical, and disuch as "Denecol" Ex301 from Nagase. Sorbitol polyglycidyl ether, such as DCE-358® from Dixie Chemical, and DCE-358® from Dixie Chemical, an

The present compositions contain a functional amount of at least one active catalyst. By functional amount is meant a quantity which will permit a VOC of less than about 3.0 and which minimizes cure volatiles, eg., acetic acid. While a blend of two or more catalysts can be used in the present compositions, at least one catalyst in the blend should be below the cure temperature, that is, at least about 15°C below the normal curing temperature. Particularly benactive below the cure temperature, that is, at least about 15°C below the normal curing temperature. Particularly benactive below the present invention are tertiary amine catalysts such as triethylene diamine, Bis(2-dimethyl aminoethyl)ether and N,N,N1,N1-tetramethylethylenediamine.

The active catalysts can be used alone or in combination with one or more additional catalysts, such as onium compounds including quaternary phosphonium and quaternary ammonium. Examples of phosphonium catalysts which can be used in catalysts blends in accordance with the present invention are benzyl triphenyl phosphonium chloride; ethyl triphenyl phosphonium bromide; tetra butyl phosphonium chloride; tetra butyl phosphonium bromide; tetra butyl phosphonium iodide; benzyl triphenyl phosp

The compositions of the present invention further comprise about from 0.01 to 5 wt %, by weight of the binder components (a) and (b), of at least one surface tension reducing agent in an amount sufficient to wet the surface onto which it is applied. A wide variety of surfactants can be used, depending on the particular coating formulation and the surface onto which it is applied. However, conventional silicone and fluorocarbon surfactants have been found to be particularly satisfactory.

The coating compositions of the present invention are formulated into high solids coating systems dissolved in at least one solvent. The solvent is usually organic. Preferred solvents include aromatic hydrocarbons such as petroleum naphtha or xylenes; ketones such as methyl amyl ketone, methyl isobutyl ketone, methyl ethyl ketone or acetone;

esters such as butyl acetate or hexyl acetate; and glycol ether esters such as propylene glycol monomethyl ether

The coating compositions of the present invention can also contain conventional additives such as pigments, stabilizers, rheology control agents, flow agents, toughening agents and fillers. Such additional additives will, of course, depend on the intended use of the coating composition. Fillers, pigments, and other additives that would adversely effect the clarity of the cured coating will not be included if the composition is intended as a clear coating.

The compositions of the present invention have a volatile organic content of less than about 360g of organic solvent per litre of curable composition (3.0 pounds per gallon), that is, at least about 65 wt % solids. The coating compositions are typically applied to a substrate by conventional techniques such as spraying, electrostatic spraying, roller coating, dipping or brushing. The present formulations are particularly useful as a clear coating for outdoor articles, such as automobile and other vehicle body parts. The substrate is generally prepared with a primer and or a color coat or other surface preparation prior to coating with the present compositions. The present coating compositions can be applied using conventional techniques such as wet-on-wet applications over solvent borne basecoats, or over dried water borne basecoats. The ability to apply the present compositions by spraying techniques with the unusually low VOC

After application to a substrate, the present compositions are cured by heating to a temperature of about from 125 content is surprising.

to 140° C for a period of about from 15 to 90 minutes. The performance characteristics of the final cured coating composition are excellent, providing a combination of excellent gloss and durability to abrasion, sunlight and acidic rain. At the same time, the compositions provide ease of handling, resulting from all the components being present in a single formulation, good shelf life and low volatile

The present invention is further illustrated by the following specific examples, in which parts and percentages are organic content. by weight unless otherwise indicated.

#### EXAMPLE 1 25

5

15

20

35

40

50

55

A curable coating composition was prepared from an anhydride resin, co-reactant oligomeric epoxy resin, active catalyst and surface tension reducing agent.

#### (a) Anhydride Resin 30

The anhydride resin was prepared from a tetra-functional half-acid ester. The following constituents were charged to a reaction vessel equipped with a heating mantle, reflux condenser, thermometer, nitrogen inlet, and stirrer:

Portion 1	Parts by Weight
Pentaerythritol Methyl hexahydrophthalic anhydride Triethylamine	478.0 2250.0 0.5
Portion 2	Parts by Weight
Xylol (135-145C) Total	2250.0 4978.5
IOIai	

Portion 1 was charged into the reaction vessel, heated to 180°C under a nitrogen blanket and held for 30 minutes. After the hold period, the reaction mixture was cooled and Portion 2 added. 45

The solution prepared above was used to make a linear pendant anhydride. The solution was charged into a 5L flask equipped with a stirrer and a gas inlet tube, the gas inlet tube was attached to a ketene generator similar to the one described by Williams et al. in the Journal of Organic Chemistry 5,122, 1940. Ketene is bubbled through the solution until all of the acid groups have been converted to anhydride groups. Reaction progress was monitored via FTIR. Solvent was then removed under vacuum to give a linear pendant anhydride with the following characteristics:

% weight solids: 78.0 Anhydride eq. wt: 329 + /-4 (on solution basis) Acid eq. wt: 6176 + /- 1323 (on solution basis)

The resulting linear pendant anhydride was combined with active catalysts and surface tension reducing agent. Based on 100 parts by weight of the final coating formulation, 55.85 parts of the anhydride resin was first combined

with active catalysts and surface tension reducing agent, along with hindered amine light stabilizer and UV screener. A blend of two catalysts was used, including a tertiary amine catalyst available from Union Carbide as NIAX A99 (1.00 parts) and 1.44 parts of a 25% solution of tetrabutyl phosponium chloride in propylene glycol monomethylether

The surfactant added to this formulation was 6.00 parts of a 5% solution in PM acetate of a polysiloxane copolymer acetate (PM acetate). commercially available as BYK-301. Also added to this linear pendant anhydride formulation were 1.00 parts of the hindered amine light stabilizer commercially available from Ciba Giegy as Tinuvin 292 and 1.40 parts of UV screener

commercially available from Ciba Giegy as Tinuvin 384. A second formulation contained the oligomer having epoxy functionality. The material used as 24.05 parts of a diglycidyl ester commercially available from DOW as XU-71950. This was combined with 3.53 parts of butyl acetate solvent and 5.73 parts of a rheology control agent. The rheology control agent was silica dispersed in epoxy. The silica dispersion consisted of 38.5 parts of the same diglycidyl ester and 51.5 parts of PM acetate, to which was added 10 parts of a hydrophobic silica commercially available from Degussa as

Aerosil R-972®. The silica dispersion was well mixed and subjected to grinding in a sand mill.

The two formulations, respectively containing the linear pendant anhydride and oligomer having epoxy functionality were combined in the ratios shown and sprayed onto primed metal panels coated with a basecoat and cured at 255°F.

The cured coating exhibited outstanding appearance and durability.

#### Claims 20

30

40

50

55

-/--

- 1. A curable coating composition comprising organic solvent and binder comprising:
- (a) an anhydride resin having a weight average molecular weight of less than about 2000 that contains a central moiety, and an average of at least three pendant, terminal, non-cyclic anhydride moieties bonded to 25
  - (b) an oligomer having epoxy functionality of at least 2 and having a weight average molecular weight of less
  - (c) a functional amount of at least one active tertiary amine catalyst, and than about 1500;
  - (d) 0.01 to 5 %, by weight of the binder components (a) and (b) together, of a surface tension reducing agent in an amount sufficient to allow the coating composition to wet the surface onto which it is applied, the composition having a volatile organic content of less than about 360g/l (3.0 pounds per gallon), and wherein the ratio of equivalents of epoxy to anhydride is about from 0.7 to 1.4.
- 2. A composition as claimed in Claim 1 comprising an additional catalyst selected from at least one phosphonium 35 compound.
  - 3. A composition as claimed in Claim 2 wherein the phosphonium compound comprises a tetrabutylphosphonium
  - A composition as claimed in Claim 1, Claim 2 or Claim 3 wherein the anhydride resin consists essentially of the reaction product of pentaerythritol, a methyl hexahydrophthalic anhydride and ketene.
- 5. A composition as claimed in Claim 1, Claim 2 or Claim 3 wherein the anhydride resin has from 3 to 4 pendant, non-cyclic anhydride moieties bonded to each central moiety. 45
  - 6. A curable coating composition as claimed in any one of Claims 1 to 5 wherein the oligomer is selected from glycidyl ethers of low molecular weight polyols, polyglycidyl esters of polyacides, glycidyl methacrylate or glycidyl acrylate containing acrylic polymers or a compatible mixture of any of the above.
  - 7. A composition as claimed in Claim 6 wherein the oligomer consists essentially diglycidyl ester of 1,2-cyclohexane dicarboxylic acid.
  - A curable coating composition as claimed in any one of Claims 1 to 7 applied to a substrate.
  - 9. A method for making a heat-curable composition in an organic solvent comprising controlling (a) the amount of volatiles produced during cure and (b) the surface tension of the coating by admixing:

- i) an anhydride resin having a weight average molecular weight less than about 2000 that contains a central moiety and an average of at least three pendant, terminal, non-cyclic anhydride moieties bonded to each
- ii) an oligomer having epoxy functionality of at least 2 and a weight average molecular weight less than about
- iii) a functional amount of at least one active catalyst selected from tertiary amine catalyst; and
- iv) 0.01 to 5 percent by weight of (i) plus (ii) of a surface tension reducing agent;
- the method also comprising employing the catalyst in step (iii) to control the amount of volatiles during cure; and employing the surface tension reducing agent in step (iv) to control the surface tension of the coating. 10

5

20

25

30

40

- 1. Härtbare Beschichtungszusammensetzung, umfassend organisches Lösungsmittel und Bindemittel, umfassend: 15
  - (a) ein Anhydridharz mit einem Gewichtsmittel des Molekulargewichts von weniger als ungefähr 2.000, welches eine Zentralgruppierung und einen Durchschnitt von mindestens drei endständigen, nicht-cyclischen Anhydridseitengruppierungen, die an jede Zentralgruppierung gebunden sind, enthält;
  - (b) ein Oligomer mit einer Epoxyfunktionalität von mindestens 2 und einem Gewichtsmittel des Molekulargewichts von weniger als ungefähr 1.500;
  - (c) eine funktionelle Menge an mindestens einem aktiven tertiären Amin-Katalysator; und
  - (d) 0,01 bis 5 %, bezogen auf das gemeinsame Gewicht der Bindemittelkomponenten (a) und (b), eines die Oberflächenspannung senkenden Mittels in einer ausreichenden Menge, die es ermöglicht, daß die Beschichtungszusammensetzung die Oberfläche, auf die sie aufgetragen wird, benetzt, wobei die Zusammensetzung einen Gehalt an flüchtigem organischen Material von weniger als ungefähr 360 g/l (3,0 Pounds pro Gallone) aufweist, und wobei das Verhältnis von Epoxy zu Anhydrid ungefähr 0,7 bis 1,4 beträgt.
  - 2. Zusammensetzung gemäß Anspruch 1, umfassend einen zusätzlichen Katalysator, gewählt aus mindestens einer Phosphoniumverbindung.
  - Zusammensetzung gemäß Anspruch 2, worin die Phosphoniumverbindung ein Tetrabutylphosphoniumhalogenid 35
    - 4. Zusammensetzung gemäß Anspruch 1, Anspruch 2 oder Anspruch 3, wobei das Anhydridharz im wesentlichen aus dem Reaktionsprodukt von Pentaerythritol, einem Methylhexahydrophthalsäureanhydrid und Keten besteht.
    - 5. Zusammensetzung gemäß Anspruch 1, Anspruch 2 oder Anspruch 3, wobei das Anhydridharz 3 bis 4 nicht-cyclische Anhydridseitengruppierungen, die an jede Zentralgruppierung gebunden sind, aufweist.
    - 6. Härtbare Beschichtungszusammensetzung gemäß mindestens einem der Ansprüche 1 bis 5, wobei das Oligomer aus Glycidylethem von Polyolen mit niederem Molekulargewicht, Polyglycidylestem von Polysäuren, Glycidylmethacrylat oder Glycidylacrylat, enthaltend Acrylpolymere, oder einer kompatiblen Mischung beliebiger der oben 45 genannten gewählt wird.
    - Zusammensetzung gemäß Anspruch 6, wobei das Oligomer im wesentlichen aus Diglycidylester von 1,2-Cyclo-7. hexandicarbonsäure besteht. 50
      - 8. Härtbare Beschichtungszusammensetzung gemäß mindestens einem der Ansprüche 1 bis 7, aufgetragen auf
    - Verfahren zur Herstellung einer wärmehärtbaren Zusammensetzung in einem organischen Lösungsmittel, umfassend das Regulieren (a) der Menge an während der Härtung erzeugten flüchtigen Materialien und (b) der Ober-55 flächenspannung der Beschichtung beim Vermischen:

- i) eines Anhydridharzes mit einem Gewichtsmittel des Molekulargewichts von weniger als ungefähr 2.000, welches eine Zentralgruppierung und einen Durchschnitt von mindestens drei endständigen, nichtcyclischen Anhydridseitengruppierungen, die an jede Zentralgruppierung gebunden sind, enthält;
- ii) eines Oligomers mit einer Epoxyfunktionalität von mindestens 2 und einem Gewichtsmittel des Molekulargewichts von weniger als ungefähr 1.500; 5
  - iii)einer funktionellen Menge an mindestens einem aktiven Katalysator, der aus tertiären Amin-Katalysatoren gewählt wird; und
  - iv) 0,01 bis 5 %, bezogen auf das Gewicht von (i) plus (ii), eines die Oberflächenspannung senkenden Mittels;
- wobei das Verfahren ebenfalls das Anwenden des Katalysators in Schritt (iii) zur Regulierung der Menge an flüchtigen Materialien während der Härtung und das Anwenden des die Oberflächenspannung senkenden Mittels in Schritt (iv) zur Regulierung der Oberflächenspannung der Beschichtung umfaßt. 15

50

- 1. Composition de revêtement durcissable comprenant un solvant organique et un liant, comprenant : 20
  - (a) une résine anhydride ayant un poids moléculaire moyen en poids inférieur à environ 2000, qui contient une partie centrale, et une moyenne d'au moins trois parties anhydride non cycliques, terminales, pendantes,
- (b) un oligomère ayant une fonctionnalité époxy d'au moins 2 et ayant un poids moléculaire moyen en poids 25
- (c) une quantité fonctionnelle d'au moins un catalyseur d'amine tertiaire actif, et (d) 0,01% à 5% en poids des composants du liant (a) et (b) ensemble, d'un agent de réduction de la tension de surface en quantité suffisante pour permettre à la composition de revêtement de mouiller la surface sur laquelle elle est appliquée, la composition ayant une teneur en composés volatils organiques inférieure à environ 360 g/l (3,0 livres par gallon), et dans laquelle le rapport des équivalents d'époxy à l'anhydride est 30
- 2. Composition suivant la revendication 1, comprenant un catalyseur supplémentaire sélectionné parmi au moins un 35
  - 3. Composition suivant la revendication 2, dans laquelle le composé de phosphonium comprend un halogénure de
- Composition suivant la revendication 1, la revendication 2 ou la revendication 3, dans laquelle la résine anhydride est composée essentiellement du produit de réaction du pentaérythritol, d'un anhydride méthylhexahydrophtalique 40
- 5. Composition suivant la revendication 1, la revendication 2 ou la revendication 3, dans laquelle la résine anhydride possède de 3 à 4 parties anhydride non cycliques pendantes liées à chaque partie centrale. 45
  - 6. Composition de revêtement durcissable suivant l'une quelconque des revendications 1 à 5, dans laquelle l'oligomère est sélectionné parmi des éthers de glycidyle de polyols de faible poids moléculaire, des esters de polyglycidyle de polyacides, du méthacrylate de glycidyle ou des polymères acryliques contenant de l'acrylate de glycidyle ou un mélange compatible d'un quelconque des composés ci-dessus.
    - Composition suivant la revendication 6, dans laquelle l'oligomère est essentiellement composé d'un ester de diglycidyle d'acide dicarboxylique de 1,2-cyclohexane.
- Composition de revêtement durcissable suivant l'une quelconque des revendications 1 à 7, appliquée sur un subs-55
  - Procédé de préparation d'une composition thermodurcissable dans un solvant organique, comprenant le contrôle

(a) de la quantité de composés volatils produits pendant le durcissement, et (b) de la tension de surface du revêtement, en mélangeant :

- i) une résine anhydride ayant un poids moléculaire moyen en poids inférieur à environ 2000, qui contient une partie centrale et une moyenne d'au moins 3 parties anhydride non cycliques terminales pendantes, liées à
- ii) un oligomère ayant une fonctionnalité époxy d'au moins 2 et un poids moléculaire moyen en poids inférieur
- iii) une quantité fonctionnelle d'au moins un catalyseur actif sélectionné parmi les catalyseurs amine tertiaire, et à environ 1500;
- iv) 0,01 à 5%, en poids de (i) plus (ii), d'un agent de réduction de la tension de surface;

5

10

15

20

25

30

35

40

45

50

55

le procédé comprenant également l'emploi du catalyseur à l'étape (iii) pour contrôler la quantité de composés volatils pendant le durcissement, et l'emploi de l'agent de réduction de la tension de surface à l'étape (iv) pour contrôler la tension de surface du revêtement.